

Claim Amendments

1. **(currently amended)** A pharmaceutical composition for intramammary administration to a non-human mammal, **wherein:**

the composition comprises: comprising

an antibacterial agent,

prednisolone, and

a pharmaceutically acceptable carrier; **and , wherein**

the composition comprises at least 20 mg of prednisolone **per** **[[/]]** unit dose.

2. **(currently amended)** The composition according to claim 1, **comprising wherein** the **composition comprises** prednisolone in an amount of 20 to 40 mg **per** **[[/]]** unit dose.

3. **(currently amended)** The composition according to claim 2, **comprising wherein** the **composition comprises** prednisolone in an amount of 20 to 30 mg **per** **[[/]]** unit dose.

4. **(previously presented)** The composition according to claim 1, wherein the antibacterial agent is a cephalosporin.

5. **(previously presented)** The composition according to claim 4, wherein the cephalosporin is cephapirin.

6. **(previously presented)** The composition according to claim 4, wherein the cephalosporin is cefquinome.

7. **(currently amended)** The composition according to claim 1, **wherein the composition comprises comprising** the antibacterial agent in an amount of 10 to 500 mg **per** **[[/]]** unit dose.

8. **(withdrawn)** A process for preparing a pharmaceutical composition according to claim 1, comprising the steps of mixing an oil and one or more pharmaceutically acceptable additives to form a carrier, and suspending the antibacterial agent and the prednisolone in the carrier.

9. **(Canceled).**